

# RDX Remote Raman Detection on NATO SET-237 Samples

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## ABSTRACT

Ultraviolet Raman spectroscopy measurements have been performed at DLR Lampoldshausen to detect NATO SET-237 standard samples of RDX. The main goal was to quantify the minimum requirements for an unambiguous identification in remote detection (60 cm distance) with a commercial Czerny-Turner spectrometer coupled with a CCD camera. Well-defined distribution of explosives on surfaces were tested as standardized samples. Therefore, Raman spectra of RDX have been acquired for different sample concentrations (50, 250, and 1000  $\mu\text{g}/\text{cm}^2$  respectively) and under several laser energies (1.5, 3.0 and 5.0 mJ/pulse respectively) at 355 nm excitation wavelength. The lowest possible reproducible surface concentration (50  $\mu\text{g}/\text{cm}^2$ ) was detected with excitation energy of 3 mJ/pulse in the described configuration.

Keywords: Raman spectroscopy, remote detection, explosives, RDX, laser diagnostics, standard samples

## 1. INTRODUCTION

The use of explosives and precursor materials is increasing to intentionally harm people and damage properties. In order to prevent attacks, a reliable detection and identification system is needed. However, the detection of these materials and their precursors remains a challenge. Often, only traces of hazardous materials are available to prevent an immediate threat. Hence there is a high demand on the sensitivity of the detection technologies. At the same time, a high specificity is necessary in order to discriminate an explosive from an interfering background or harmless substances. Therefore, obtaining a well-defined spectral signature that leads to a clear identification and a low rate of false alarm is very difficult from large molecules like explosives; moreover, the background material might cover, absorb or distort the signal causing more uncertainty. Different types of technology should be employed depending on the type of hazardous compound and environmental conditions. In addition, it is desirable to achieve the widest possible range of detectable hazardous substances and, at the same time, to reduce the acquisition time and laser energy. For these reasons, a comprehensive investigation of the detection possibilities and their optimization for the various warfare agents is mandatory [1]. Among all the available technologies [1,2], the ones with the best potential of being fast and able to detect and identify many explosive compounds and upgradable to new threats are laser-based spectroscopic methods for trace detection. In this context, the presented work is focused on short-pulse Raman spectroscopy chosen for remote detection. Raman spectroscopy was selected for its unique capability of being fast-responsive, highly sensitive, selective and able to identify different explosive compounds [3,4] / chemical threats [5]. The final goal is to find the minimal detectable amount of the target compound ( $\mu\text{g}/\text{cm}^2$ ) and minimal energy density ( $\text{mJ}/\text{cm}^2$ ) required for a clear identification and (in a future step) classification. For this purpose, several samples of perhydro-1,3,5-trinitro-1,3,5-triazine (RDX) [6] were tested. The preparation of the samples adds an additional challenge: different methods such as dropping and drying (drop casting) or spray deposition present an easy and fast procedure [1]; however, both methods are not the most recommended choice in term of reproducibility of the sample type since they provide inhomogeneous distributions (coffee ring pattern) [7]. Among these possibilities, drop-on-demand (DOD) inkjet printing is becoming a promising method for producing standardized test materials with high precision, reproducibility [8] and accuracy [9] especially at very low concentrations. Hence all the RDX samples here presented were prepared in different concentrations as NATO SET-237 [10] standards using DOD inkjet printing method developed by the Fraunhofer Institute of Chemical Technology (ICT) [11]. The thresholds and limits of selectivity and sensitivity are recorded and discussed, as well as the associated laser energy and the detection distance (fixed at 60 cm in this work) for the respective explosive and background.

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## 2. EXPERIMENTAL SETUP

A sketch of the Raman system setup [12] is shown on Figure 1. An injection seeded Nd:YAG laser (*Spectra Physics Quanta Ray Pro 230*) with a repetition rate of 10 Hz (pulse duration < 8 ns) was frequency doubled to produce a 532 nm laser beam; this beam was used to pump a narrowband dye laser (*Sirah Cobra-Stretch*). The green light beam left over after the doubling was internally frequency tripled to produce a 355 nm laser beam. The UV light was then sent to a second exit of the laser and frequency mixed with the output of the dye laser inside a commercial mixing module (*Sirah SFM-355*). Depending of the choice of the dye solution or dye mixture, the mixing module is able to produce UV light in the range from 200 nm to 240 nm. The choice of the output wavelength varies by experimental requirements and by the compound under examination (signal optimization). After exiting the mixing module, the UV laser beam was relayed by a series of UV mirrors first through a beam attenuator (consisting on a rotating half-wave plate and a polarizer) to control the energy (or energy density) of the laser beam, and then through a telescope (consisting of two lenses of 150 mm and 75 mm focal length, FL) to reduce by half the beam size and to collimate it to the sample. Finally, the 8 mm diameter collimated UV laser beam was sent to the sample to excite the Raman transition. It can be noticed that the beam is not focused onto the sample in order to do not destroy or damage it and to increase the sampling area.

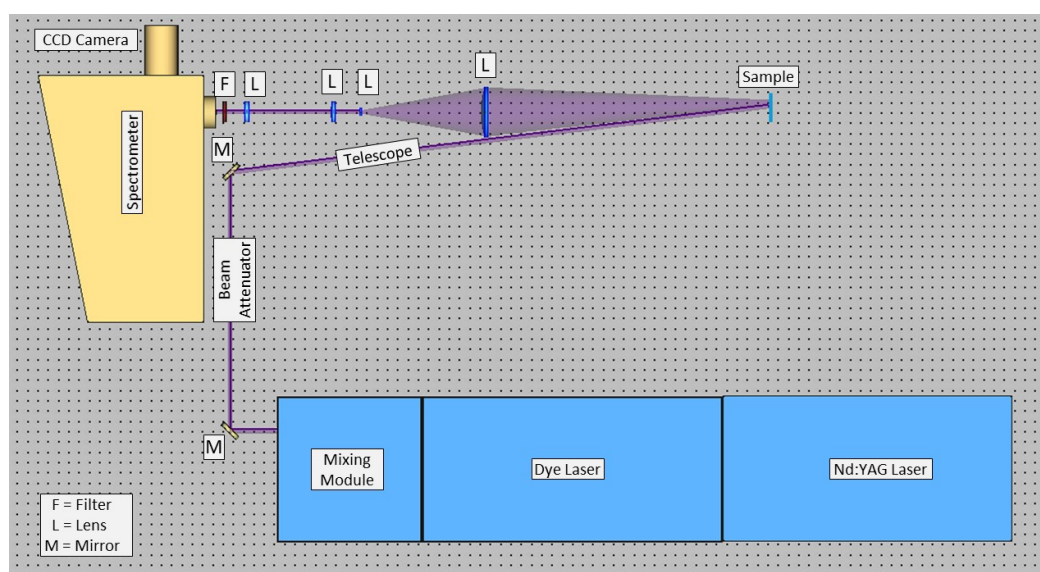


Figure 1. Sketch of the Raman optical setup.

The generated Raman signal was collected by a 75 mm diameter, 200 mm FL lens placed at a distance of about 60 cm from the sample. Subsequently, a couple of lenses collimated the signal which is finally focused on a 200  $\mu\text{m}$  Czerny-Turner spectrometer's (*Horiba iHR550*) slit by a 75 mm FL lens. A blocking UV filter (blocking wavelength depended on the exiting beam) was placed in front of the slit in order to eliminates residual scattered laser light. Inside the spectrometer, the Raman signal was first dispersed by a 2400 groves/mm holographic grating (blaze 250 nm, resolution 0.06 nm) and then focused on a couple charged device (CCD) *Horiba Symphony II*. The camera (2048 x 512 pixel array, maximum resolution 0.01 nm) was vertically binned in order to collect a spectrum from the Raman signal. The overall resolution of the system for the acquired spectra was  $15\text{ cm}^{-1}$ . Acquisition time varied form a minimum of 1 s to a maximum of 30 s depending on the sample type, concentration, distance, and laser energy.

## 1. EXPLOSIVE SAMPLES

The laboratory samples presented in this study were produced as NATO SET-237 [10] by the Fraunhofer ICT using a *GeSiM 2.1 Nano-Plotter* (~40-500 pico-liters per droplet) DOD inkjet printer [11]. Since the printing method is completely contactless, extremely small droplets of concentrated material can be homogeneously spread over a wide selection of solid backgrounds in different patterns. The requirement is a homogeneous coating of the substrate with a precisely predefined concentration and a uniform particle size. The solvent in use for the samples under examination were acetonitrile (ACN) and methyl ethyl ketone (MEK). The threat concentration (and droplet volume) varies from 10 to 30 mg/ml and it was determined by means of stroboscope observation with optical sensor as well as by a flow sensor. The plotting speed is determined by the analyte itself and its concentration into the solvent as well as the number of

droplets per point and the number of layers printed. Tables 1 summarizes the characteristics of the produced samples and the different printing parameters chosen for RDX.

Table 1. RDX samples characteristics.

RDX Sample #	Solvent	Concentration [mg/ml]	Array	Droplets per point	Layers	Final distribution [ $\mu\text{g}/\text{cm}^2$ ]
832	ACN	10	40 x 44	2	4	50
835	MEK	20	34 x 36	6	5	250
838	MEK	20	34 x 35	4	10	1000

At the beginning of a plotting process droplets of the solution are formed. By evaporation of the solvent these droplets shrink and finally solid residues remain on the substrate. Finally, the second layer is printed again and the procedure is repeated till the desired final distribution is obtained. The first NATO SET-237 of RDX samples are shown in Figure 2. In these samples preparation the explosives were printed on an area of 1 x 1 cm over an aluminum substrate.

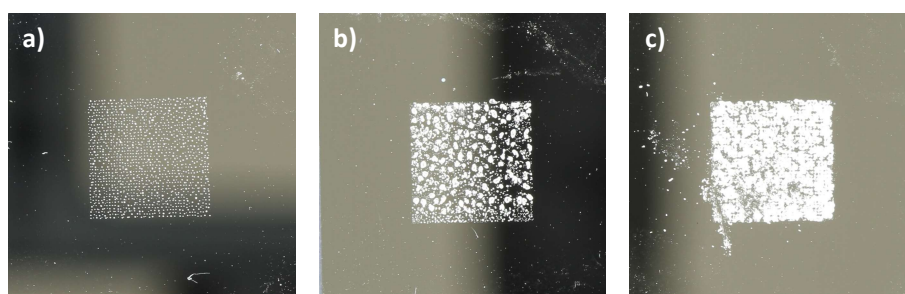


Figure 2. Macro pictures of RDX NATO-237 samples a) #832 ( $50 \mu\text{g}/\text{cm}^2$ ); b) #835 ( $250 \mu\text{g}/\text{cm}^2$ ); c) #838 ( $1000 \mu\text{g}/\text{cm}^2$ )

## 2. SAMPLES' ANALYSIS

All the samples were first analyzed to verify the compound distribution and physical properties. A preliminary analysis using the microscope *Krüss Optronic MSZ 5400* was done to determine the overall distribution on the sample over the background surface. An *Edinburgh Instrument Spectrofluorometer FS5* was employed to measure absorbance of all tested materials. This information was useful to select the best laser wavelength to maximize the Raman signal. Figure 3 compares the samples of RDX in different concentrations using the - *Krüss Optronic MSZ 5400* microscope -with a 40x magnification. The main goal was to verify the integrity and the uniformity of the droplet distribution of the samples.



Figure 3. 40x selected views of RDX samples a) #832 ( $50 \mu\text{g}/\text{cm}^2$ ); b) #835 ( $250 \mu\text{g}/\text{cm}^2$ ); c) #838 ( $1000 \mu\text{g}/\text{cm}^2$ )

It can be observed that obtaining a similar -distribution over the sample is quite challenging for RDX. The microscope images show different droplets size in the sample and this effect could be attributed to the high melting point temperature of RDX ( $206^\circ\text{C}$ ) [6]. Most industrial printing devices use piezo- electrically or thermally generated pressure pulses to eject droplets from a fluid cavity: during this procedure the solution solvent-explosive is heated up to  $40^\circ\text{--}50^\circ\text{C}$ . Under this condition RDX is still in solid form, hence its conglomerate may result in partial droplet released. In addition, looking from Figure 3a to 3c, it can be inferred that at higher overall explosive distribution it is not possible to achieve uniformity of the sample. The images show a more uneven distribution of particles and uneven size of the droplets with increasing the amount of the deposited material. The multi-layer approach to produce higher concentration is unable to provide a uniform distribution and the particles of RDX tend to cluster together. The subsequent evaporation of the

solvent has the only effect to spread the material over the background surface creating local inhomogeneous patterns. However, all the samples were well confined into the 1 cm<sup>2</sup> and the achieved distribution, although non-fully-uniform, was found much better than the one obtained by the drop-cast method [7].

The *Edinburgh Instrument Spectrofluorometer FS5* was employed to obtain the absorbance spectrum of the samples. The explosive was tested using the compound dissolved into a solution. Concentrated solution of RDX in ACN were prepared and tested inside a cuvette. A cuvette filled with pure ACN was tested as reference background. The concentration was experimentally found based on the strength of the transmission spectrum. Exceeding an imposed value implied saturating the spectrometer (hence damaging it). Concentration of RDX was set to  $5.44 \times 10^{-3}$  mg/ml ( $2.45 \times 10^{-5}$  mol/l). Transmittance was measured inside the Spectrofluorometer scanning each wavelength produced by a Xenon lamp from 215 to 900 nm with 1 nm step. At each wavelength, the light illuminated the sample for 0.5 s. From the measured transmittance it was possible to calculate the absorbance. Figure 4 shows the absorbance spectrum of RDX. The absorbance peak of RDX is at 215 nm (deep UV). Moving towards the visible wavelengths the absorbance drops fast and almost no light is absorbed after 380 nm. The values are also in agreement with Lu et al. [13] However, despite the peaks of the absorbance are a good indication about the strength of the Raman signal, they are also providing the strongest fluorescence. Hence, in order to avoid strong interfering fluorescence from the sample, laser sources in the range of 355 nm could be a good compromise between strength of the Raman signal and low fluorescence.

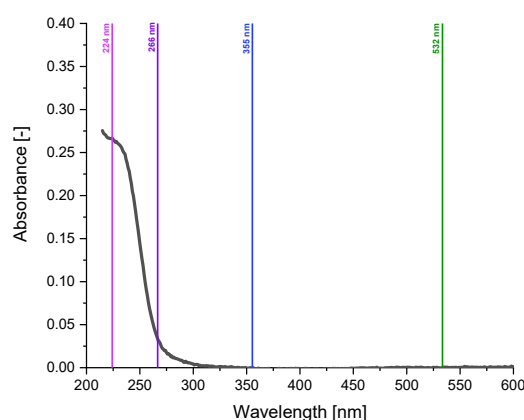


Figure 4. Absorbance spectrum of RDX. Vertical lines represent the frequently used Raman excitation wavelengths. Results for 355 nm are presented.

### 3. RESULTS AND DISCUSSION

Raman measurements were performed using different concentrations of explosive samples (50, 250 and 1000 µg/cm<sup>2</sup> respectively) and different energies (1.5, 3 and 5 mJ/pulse) of the exciting laser (355 nm wavelength at 10 Hz repetition rate). The exciting laser beam was collimated to 8 mm diameter yielding an energy density of 30, 60 and 100 mW/cm<sup>2</sup> respectively. The Raman signal was collected at a distance of about 60 cm from the target sample inside a spectrometer coupled with a CCD camera with an acquisition time of 30 s. Samples #832, #835 and #838 were chosen as representative of 50, 250 and 1000 µg/cm<sup>2</sup> of RDX concentration respectively. Figure 5 shows the Raman spectra of RDX #832 (50 µg/cm<sup>2</sup>) tested at different laser energies. The black line represents the raw data, while the thick red line represents a smoothing function consisting of an average of  $\pm 5$  adjacent points. The most known peaks [7] of the compound are marked in red.

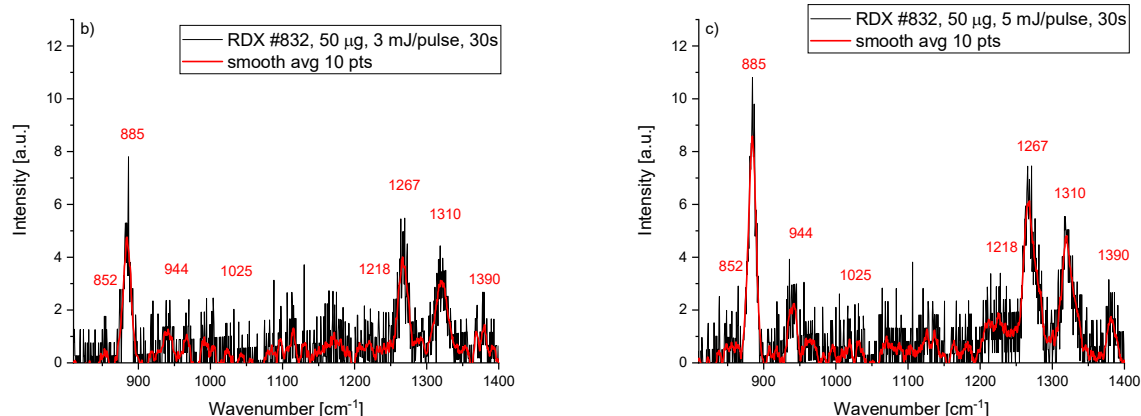
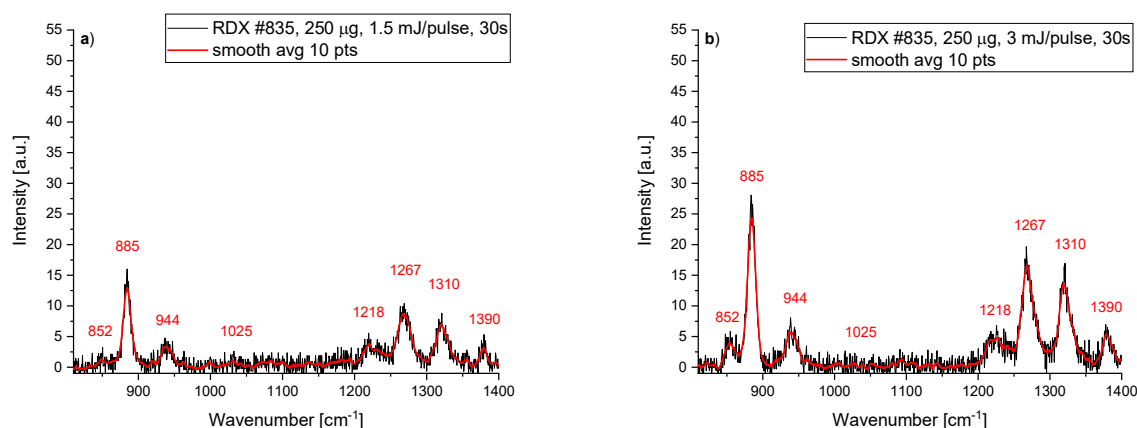


Figure 5. Raman spectra of 50  $\mu\text{g}/\text{cm}^2$  of RDX; b) laser energy 3 mJ/pulse; c) laser energy 5 mJ/pulse

It can be noticed that the Raman signature of RDX is evident in both plots. The main peak at 885  $\text{cm}^{-1}$  (mainly CN stretch) can be recognized by the background level as well as the peaks at 1267 (NN stretching and ONO stretching, maybe with  $\text{CH}_2$  twist) and 1310  $\text{cm}^{-1}$  ( $\text{CH}_2$  wag or combination) [14,15]. However, at lower laser energy, all the other spectral features are not distinguishable from the background even with the help of the smoothing function (red line). This suggests that 3 mJ/pulse is still not enough to have a clear spectral signature and more energy or higher concentration of the hazardous compound are necessary for a better identification. For this reason, the test with laser energy of just 1.5 mJ/pulse is not shown: even the main peak at 885  $\text{cm}^{-1}$  was not strong enough to be discriminated from the background noise. On the other hand, Figure 7c shows much more structured features. Despite the low concentration of RDX, peaks at 944 and 1218  $\text{cm}^{-1}$  are clearly visible, in addition to the three already mentioned main peaks. Having more information about the spectral shapes leads to a better identification of the hazardous compound and reduces the rate of false alarms. Hence, laser energy of at least 5 mJ/pulse is recommended to detect 50  $\mu\text{g}/\text{cm}^2$  (or more) RDX in the presented configuration. Figure 6 shows the Raman spectra of RDX #835 (250  $\mu\text{g}/\text{cm}^2$ ) tested at different laser energies.



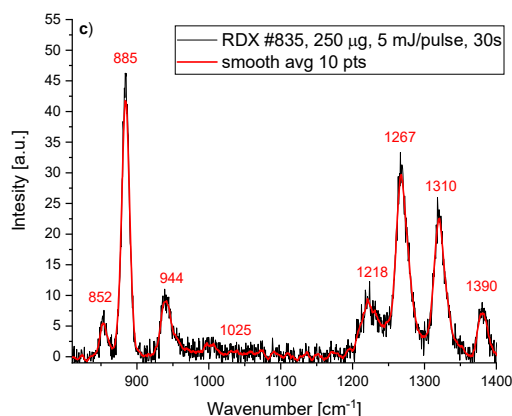
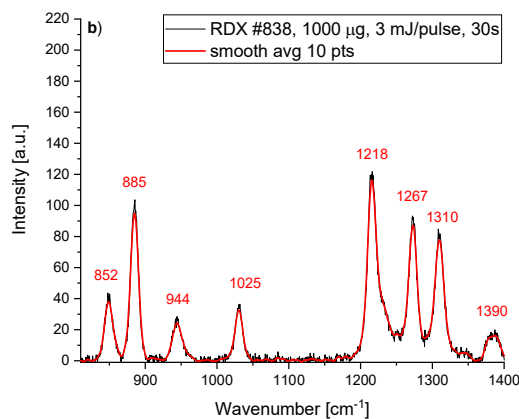
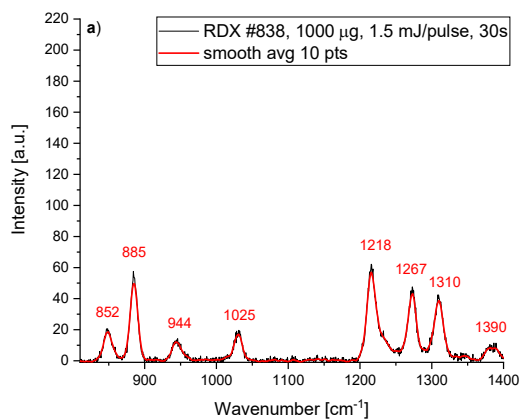


Figure 6. Raman spectra of 250  $\mu\text{g}/\text{cm}^2$  of RDX; a) laser energy 1.5 mJ/pulse; b) laser energy 3 mJ/pulse; c) laser energy 5 mJ/pulse

Again, all the strongest peak [7] of the RDX at 885, 1267 and 1310  $\text{cm}^{-1}$  [14,15] respectively are clearly distinguishable from the background in all the three spectra. In addition, peaks at 944 and 1218  $\text{cm}^{-1}$  are strong enough for identification even at 1.5 mJ/pulse energy. This implies that low laser energy is required once the concentration of the RDX is at least 250  $\mu\text{g}/\text{cm}^2$ . Of course, when more laser energy is used, the spectral features are clearer since the signal to noise ratio is much higher, as it can be noticed in Figure 8b and 8c where peaks at 852 (NN stretch plus  $\text{NO}_2$  axial scissoring) and 1218  $\text{cm}^{-1}$  (NC stretching) [14,15] are stronger in intensity. Increasing the laser energy up to 5 mJ/pulse makes the peak at 1025  $\text{cm}^{-1}$  (NC stretch with some  $\text{CH}_2$  rocking) [14,15] visible, even though is not completely discriminated by the background noise. In conclusion, the Raman system is sensitive enough to detect at least 250  $\mu\text{g}/\text{cm}^2$  of RDX using just 1.5 mJ/pulse laser energy. Figure 7 shows the Raman spectra of RDX #838 (1000  $\mu\text{g}/\text{cm}^2$ ) tested at different laser energies.





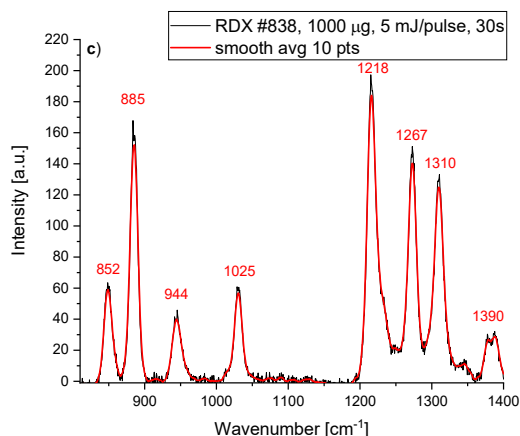


Figure 7. Raman spectra of 1000  $\mu\text{g}/\text{cm}^2$  of RDX; a) laser energy 1.5 mJ/pulse; b) laser energy 3 mJ/pulse; c) laser energy 5 mJ/pulse

In this last case, RDX is concentrated enough to be detected at each tested laser energy. The signal to noise ratio is high in all the three pictures. Interesting that at these conditions the peak at 1025  $\text{cm}^{-1}$  can be used as an additional discrimination factor to identify the compound. An additional peak is also visible at 1390  $\text{cm}^{-1}$  ( $\text{CH}_2$  twisting) [14,15]. Hence at a concentration of at least 1000  $\mu\text{g}/\text{cm}^2$ , the Raman system required 1.5 mJ/pulse to be sensitive enough to clearly identify RDX at 60 cm distance. As shown in Section 4, the sample material to be examined was non-homogeneously distributed on the substrate surface. For the NATO-Printed Standard samples their nominal density value is given in  $\mu\text{g}/\text{cm}^2$ . RDX was distributed more or less evenly over the substrate surface, depending on the efficiency of the printing process on the 1 x 1  $\text{cm}^2$  sample area. As a first approximation, the signal depends on the area of the sample that interacts with the incident excitation light as shown in Figure 8. The effective sampled area was below the total available area (1  $\text{cm}^2$ ) because the incident laser beam was smaller (0.5  $\text{cm}^2$ ).

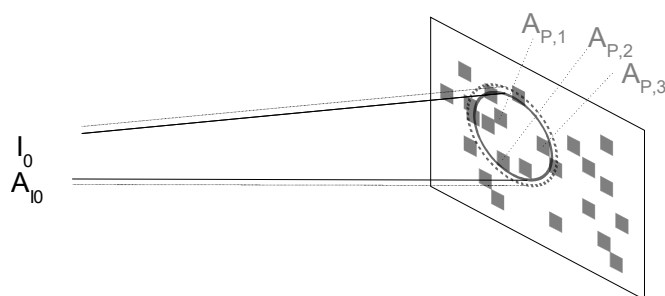


Figure 8. Sketch of the area ratios on a non-homogeneous RDX printed sample. Intensity of the exciting laser  $I_0$ , area of the laser on the sample  $A_{I0}$ , area of the individual sample particles on the background  $A_{p,i}$ ;

In the case of resonance Raman scattering (and fluorescence), absorption effects must also be considered. For RDX the penetration depth is specified in the literature [16] to be  $\sim 1.1 \mu\text{m}$ . Taking this factor into consideration, the nominal explosive density value (specified for the NATO given samples in  $\mu\text{g}/\text{cm}^2$ ) is only a limited measure for specifying the sensitivities of the measurement process. Practically, only the illuminated surface and a minimal part of the thickness contributes to the Raman signal while all the material found below the penetration depth is not detected. Using the images from the microscope analysis it was possible to compute the effective detectable quantity of explosives taking also into consideration that the area of the incident laser beam (0.5  $\text{cm}^2$ ) was smaller than the sample size. Figure 9 shows the dependency of the Raman signals of RDX as a function of the effective detectable quantity in  $\mu\text{g}$ .

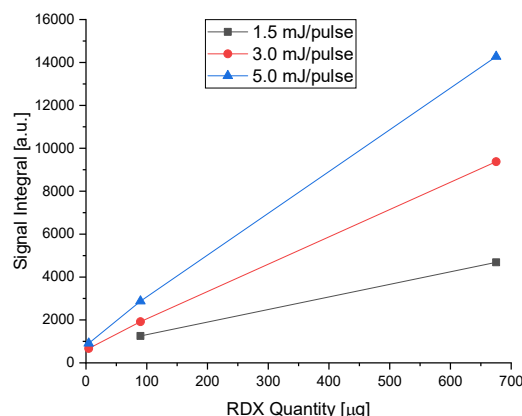


Figure 9. Detectable Raman signals of RDX as a function of effective quantity at different excitation laser pulse energy

It can be noticed that the minimal amount detectable is 4.4  $\mu\text{g}$  of RDX using at least 3 mJ/pulse energy. However, it requires a minimum of 89.7  $\mu\text{g}$  of RDX to be detected by the minimum laser energy tested (1.5 mJ/pulse).

#### 4. SUMMARY

An experimental setup for remote Raman spectroscopy with short pulse systems was optimized for the investigation of explosives on contaminated surfaces. First results were presented in this study to identify the detection limits (laser energy and compound's concentration) for RDX. The measurements were performed on printed RDX NATO SET-237 samples. The Raman signature of the explosive corresponds to the signature of the pure substance known from the literature. The minimum detectable RDX was 4.4  $\mu\text{g}$  with exciting laser energy of 3 mJ/pulse. However, the signal to noise ratio was found too low for a clear identification, so 5 mJ/pulse are suggested to avoid false alarms. Increasing further the concentration of RDX (at least 89.7  $\mu\text{g}$ ) provided always a clear identification of the explosive material even with 1.5 mJ/pulse. A further optimization of the system (pulse energies and pulse diameter, collecting optics, detection device) will be performed to possibly decrease the detectable amount limits. Moreover, the preparation of a suitable sample plays an essential role in determining the sensitivities of the technologies used and their specified standard concentration ( $\mu\text{g}/\text{cm}^2$ ) isn't a realistic measure for the effective detection sensitivity, especially in the case of inhomogeneous surfaces with thick sample particles. A better estimation of the sensitivity can be determined on homogeneous layers with a thickness equal to the penetration depth of the exciting laser light.

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